## ABSTRACT OF THE DISCLOSURE

Data presented herein provide a molecular mechanism for circadian gene mPer2 in DNA damage response and tumor suppression  $in\ vivo$ . Mice deficient in mPer2 gene display neoplastic phenotypes. These mice are deficient in p53-mediated apoptosis in thymocytes and have increased tumor occurrences after  $\gamma$ -radiation. Core circadian genes are induced by  $\gamma$ -radiation in wild-type mice but not in mPer2 mutant mice. Temporal expression of genes involved in cell cycle regulation and tumor suppression, such as c-Myc,  $Cyclin\ D1$ ,  $Cyclin\ A$ , Mdm-2 and  $Gadd45\alpha$  is dependent on  $mPER2\ in\ vivo$ .

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